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Computer-Aided Detection for Prostate Cancer Detection based on Multi-Parametric Magnetic Resonance Imaging

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Abstract—Prostate cancer (CaP) is the second most diagnosed cancer in men all over the world. In the last decades, new imaging techniques based on magnetic resonance imaging (MRI) have been developed improving diagnosis. In practice, diagnosis is affected by multiple factors such as observer variability and visibility and complexity of the lesions. In this regard, computer-aided detection and diagnosis (CAD) systems are being designed to help radiologists in their clinical practice. We propose a CAD system taking advantage of all MRI modalities (i.e., T₂-W-MRI, DCE-MRI, diffusion weighted (DW)-MRI, MRSI). The aim of this CAD system was to provide a probabilistic map of cancer location in the prostate. We extensively tested our proposed CAD using different fusion approaches to combine the features provided by each modality. The source code and the dataset have been released.

I. INTRODUCTION

Current prostate cancer (CaP) screening consists of 3 different stages. First, prostate-specific antigen (PSA) control is performed to distinguish between low- and high-risk CaP. To assert such diagnosis, samples are taken during prostate biopsy and analyzed to make an accurate prognosis of the CaP.

Although PSA screening has been shown to improve early detection of CaP [1], its lack of reliability motivates further investigations using magnetic resonance imaging (MRI)-based computer-aided detection and diagnosis (CAD). Consequently, current research is focused on identifying new biological markers to replace PSA-based screening [2]. Until such research comes to fruition, these needs can be met through active-surveillance strategy using multiparametric MRI (mp-MRI) techniques [3].

Lemaître *et al.* recently reviewed more than 50 research works that focused on CAD system for CaP [4]. These studies are based on CAD systems that consist of the following steps: (i) pre-processing, (ii) segmentation, (iii) registration, (iv) feature detection, (v) feature selection-extraction, and (vi) finally classification.

The reviewed mp-MRI-based CAD used 2 to 3 MRI modalities among T₂ Weighted (T₂-W)-MRI, dynamic contrast-enhanced (DCE)-MRI, and diffusion weighted (DW)-MRI, discarding the potential discriminative power of magnetic resonance spectroscopy imaging (MRSI). Furthermore, only half of these studies tackled the challenging detection of CaP in the central gland (CG). Additionally,

none of the works investigated the issue related to feature balancing when developing their CAD systems. Finally, none of the datasets nor source codes used have been released, making impossible the possibilities to compare the methods.

In this work, we propose a CAD system to detect CaP in peripheral zone (PZ) and CG, using the 4 aforementioned MRI modalities. The ultimate goal of the current CAD system is to provide a probabilistic map of the cancer within the prostate. Therefore, each voxel in the prostate will be classified as healthy or cancerous. The dataset used and the source code developed are released for future comparisons and reproducibility.

II. METHODOLOGY

A. Materials

The mp-MRI data are acquired from a cohort of patients with higher-than-normal level of PSA. Acquisition is achieved with a 3 T whole body MRI scanner (Siemens Magnetom Trio TIM, Erlangen, Germany) using sequences to obtain T₂-W-MRI, DCE-MRI, DW-MRI, and MRSI. In addition of the MRI examination, these patients also have undergone a transrectal ultrasound (TRUS) guided-biopsy. The dataset is composed of 17 which all have biopsies that were positive for CaP. From those 12 patients have a CaP in the PZ, 3 patients have CaP in the CG, 2 patients have invasive CaP in both the PZ and the CG. An experienced radiologist segmented the prostate organ — on T₂-W-MRI, DCE-MRI, and apparent diffusion coefficient (ADC) — as well as the prostate zones — i.e., PZ and CG —, and CaP on the T₂-W-MRI. The full description and the data set are available at I2Cvb website¹ [5].

B. CAD pipeline for CaP

Our mp-MRI CAD system consists of 7 different steps: pre-processing, segmentation, registration, feature detection, feature balancing, feature selection/extraction, and finally classification. The different source codes are publicly available².

¹<http://i2cvb.github.io/>

²<https://github.com/I2Cvb/mp-mri-prostate>

1) *Pre-processing*: Normalization is, a crucial step to reduce the inter-patient variations which allows to improve the learning during the classification stage. However, the MRI modalities provide specific type of data — static vs. dynamic information, images vs. signals — that required a dedicated pre-processing. Therefore, we pre-process differently the data: T₂-W-MRI is normalized using a Rician a-priori that has been shown to be better than the traditional z -score [6]. In contrast to T₂-W-MRI, in ADC map the probability density function (PDF) within the prostate does not follow a known distribution and thus one cannot use a parametric model to normalize these images and a non-parametric piecewise-linear normalization [7] is the best option for this case. DCE-MRI is a dynamic sequence and the data are normalized based on a mean kinetic expression registration as proposed in [5]. Finally, the MRSI modality has been pre-processed to correct the phase, suppress the baseline, and align the frequencies [8].

2) *Segmentation and registration*: For this work, our radiologist has manually segmented the prostate organs on the different modalities. However, the segmented prostate needs to be registered before to extract features. Therefore, the patients motion during the DCE-MRI is corrected using a rigid registration with an mean squared error (MSE) similarity metric and a gradient descent optimizer. Subsequently, the T₂-W-MRI and DCE-MRI are co-registered using a rigid transformation and the delineation of the prostate gland, using the same metric and optimizer previously mentioned. ADC maps and T₂-W-MRI are also co-registered with the same strategy. Additionally, volumes from all modalities have been interpolated to the resolution of T₂-W-MRI.

3) *Feature detection*: Similarly to the pre-processing, specific features are extracted depending of the specificity of each MRI modality.

T₂-W-MRI and ADC map features Additionally to the normalized intensity, edge- and texture-based features are commonly extracted from T₂-W-MRI and ADC map. The following set of filters characterizing edges have been used: (i) Kirsch, (ii) Laplacian, (iii) Prewitt, (iv) Scharr, (v) Sobel, and (vi) Gabor. Except for the Kirsch filter, the other filters are applied in 3D, taking advantage of the volume information instead of slice information, as it is usually done. Additionally, features based on phase congruency are computed [9]. To characterize the local texture, both second-order gray-level co-occurrence matrix (GLCM)-based features [10] and rotation invariant and uniform local binary pattern (LBP) [11] are extracted. To encode 3D information, the 13 first Haralick features are computed for the 13 possible directions. For the same reason, the LBP codes are computed for the three-orthogonal-planes of each MRI volume. All these features are extracted at each voxel of the volume.

DCE-MRI features In brief, the entire enhanced signal, semi-quantitative [12], and quantitative-based models [13], [14], [15], [16] are computed.

MRSI features Three different techniques are used to ex-

tract discriminative features: (i) relative quantification based on metabolite quantification, (ii) relative quantification based on bounds integration, and (iii) spectra extraction from 2 ppm to 4 ppm [5].

Anatomical features Four different metrics are computed based on the relative distance to the prostate boundary as well as the prostate center, and the relative position in the Euclidean and cylindrical coordinate systems [17], [18].

4) *Feature balancing*: Imbalanced dataset is a common problem in medical imaging. The number of cancerous voxels is much lower than the number of “healthy” voxels for a patient. This problem compromises the learning process. Solving the problem of imbalanced is equivalent to under- or over-sampling part of the dataset to obtain equal number of samples in both classes. In this regard, the imbalanced dataset was under-sampled using the different variant of nearmiss (NM) [19] and the instance-hardness-threshold (IHT) [20] algorithm. In addition, the dataset was also balanced using over-sampling methods, namely different variant of synthetic minority over-sampling techniques (SMOTE) [21], [22]. Those algorithms were developed and made publicly available in the `scikit-learn-contrib-imbalanced-learn`³ python package [23].

5) *Feature selection and extraction*: Feature selection and extraction are used in our experiment. MRSI and DCE-MRI are decomposed using three feature extraction methods: principal components analysis (PCA), sparse-PCA, and independent components analysis (ICA). Additionally to feature extraction, two methods of feature selection are used: (i) the one-way analysis of variance (ANOVA) and (ii) the Gini importance obtained while learning the random forest (RF) classifiers. The `scikit-learn`⁴ python package provides all those methods and was used in our experiment [24].

6) *Classification*: RF has been chosen as our base classifier to perform classification of individual modality as well as the combination of modalities. RF and decision trees do not require to scale features and provide feature selection by analyzing the feature importance, derived from the impurity improvement successive splits. Additionally, we use stacking to create ensemble of base learners using a meta-classifier [25], namely AdaBoost (AdB) and Gradient Boosting (GB).

III. RESULTS AND EVALUATION

Various experiments were run in order to optimize the balancing and the feature selection strategies [5]. We found that once all features are concatenated together, nearmiss-3 (NM-3) [19] is the method providing the best enhancement of the classification performance with an area under the curve (AUC) of 0.824 ± 0.076 . Therefore, with this optimal balancing, we here report the final step consisting of three strategies: (i) the selected features from each modality (i.e.,

³<http://contrib.scikit-learn.org/imbalanced-learn/>

⁴<http://scikit-learn.org/stable/>

TABLE I

SELECTED FEATURE AND NUMBER OF OCCURRENCE FOR T₂-W-MRI, ADC MAP, AND ONE ALL THE FEATURES ARE CONCATENATED.

T ₂ -W-MRI	ADC	T ₂ -W-MRI	ADC	DCE-MRI	MRSI
8 edges	1 DCT	113 Gabor filters	53 Gabor filters	14 samples	78 samples
155 Gabor filters	32 Gabor filters	1 phase congruency	2 phase congruency		
2 Haralick features	1 phase congruency	4 edges			
1 intensity		1 intensity			
4 LBP					
2 phase congruency					
172 features	34 features	267 features			

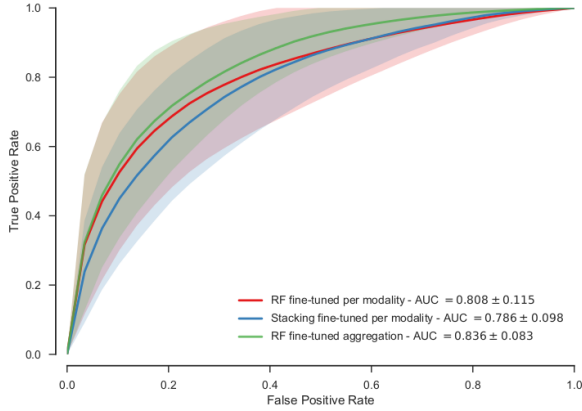


Fig. 1. Analysis of feature combination approaches after fine tuning through balancing and feature selection/extraction.

331 features) are concatenated together and used in a RF classifier, (ii) the selected features from each modality (i.e., 331 features) are used to train a stacking classifier with a GB as meta-classifier, and (iii) the selected features from the concatenated set of features (i.e., 267 features) are used to train a single RF classifier.

The selected features are presented in Table I which highlights some interesting facts regarding the most efficient features. On the one hand, the Gabor filters and the phase congruency are always selected, independently of the strategy and modality during the feature selection process. Additionally, edge filters — i.e., Kirsch, Prewitt, Scharr, and Sobel — have been only selected for the T₂-W-MRI. A possible explanation might be due to the fact that T₂-W-MRI is the modality with the highest spatial resolution and in which the level of details is the most important. Subsequently, the intensity feature of the T₂-W-MRI modality is always selected, implying that our normalization method proposed in [6] is efficient.

The experiments were performed in a leave-one-patient-out cross-validation (LOPO CV) fashion and a receiver operating characteristic (ROC) analysis is carried out. The comparative results are shown in Fig. 1. In overall, classification using the fine-tuned features improve the classification performance. The third classification configuration is, however, the one which outperforms others with an AUC of 0.836 ± 0.083 . The improvement in terms of AUC is of 0.028 and 0.050 compared with the 1st and 2nd configurations,

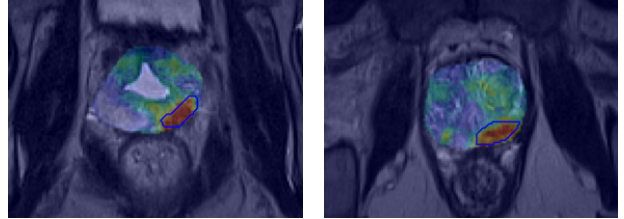


Fig. 2. Illustration the resulting detection of our mp-MRI CAD for CaP detection. The blue contours corresponds to the CaP while the jet overlay represents the probability.

respectively.

In a clinical setting, the AUC score is categorized in 3 levels: (i) “acceptable” discrimination for an AUC ranging from 0.7 to 0.8, (ii) “excellent” discrimination for an AUC ranging from 0.8 to 0.9, and “outstanding” discrimination when the AUC is over 0.9 [26]. Therefore, the combination of all MRI modalities in conjunction with fine-tuning allow to upgrade our CAD system from an “acceptable” to an “excellent” discrimination level.

To illustrate qualitatively the results of our mp-MRI CAD system, 2 diverse examples are presented in Fig. 2 by overlapping the probability map of having a CaP with the original T₂-W-MRI slice.

IV. CONCLUSION

In this paper, we presented one of the the first CAD system using all the mp-MRI modalities for prostate cancer detection. Indeed, MRSI has nearly never been used together with the other modalities. With an extensive validation approach to select the best features, the best balancing strategy as well as the best classifier, we obtained results on a rather complicated dataset of 17 patients with an average AUC of 0.836 ± 0.083 which put our system in the state-of-the-art, even so different CADs were tested on different datasets. In addition, we released the source code and the dataset allowing for reproducibility.

As avenues for future research, one could switch from voxel-based classification to super-voxel classification such that spatial structure are classified instead of voxel. In addition, the registration relies on the segmentation of the prostate gland which was provided by our doctors. To be

used in a clinical environment, this step need to be fully automatized.

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